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Hormonal contraception does not increase women's HIV acquisition risk in Zambian discordant couples, 1994–2012

Kristin M. Wall^{a,b,*}, William Kilembe^a, Bellington Vwalika^{a,d}, Naw Htee Khu^a, Ilene Brill^c, Elwyn Chomba^{a,e}, Brent A. Johnson^f, Lisa Haddad^{a,g}, Amanda Tichacek^a, and Susan Allen^a Rwanda Zambia HIV Research Group, Department of Pathology & Laboratory Medicine, School of Medicine and Hubert Department of Global Health, Rollins School of Public Health, Emory University, Atlanta, GA, USA

^bDepartment of Epidemiology, Rollins School of Public Health, Laney Graduate School, Emory University, Atlanta, GA, USA

^cDepartment of Epidemiology, Ryals School of Public Health, University of Alabama at Birmingham, Birmingham, AL, USA

^dDepartment of Gynecology and Obstetrics, School of Medicine, University of Zambia, Lusaka, Zambia

eMinistry of Community Development, Mother and Child Health, Lusaka, Zambia

^fDepartment of Biostatistics and Computational Biology, School of Medicine and Dentistry, University of Rochester Medical Center, Rochester, NY, USA

^gDepartment of Gynecology and Obstetrics, Emory University, School of Medicine, Atlanta, GA, USA

Abstract

Objective—To determine the impact of hormonal contraceptive methods on risk of HIV acquisition among HIV-negative women cohabiting with HIV-positive male partners.

Study design—From 1994–2012, HIV discordant couples recruited from a couples' voluntary HIV counseling and testing center in Lusaka, Zambia were followed longitudinally. HIV-negative partners were tested quarterly. This analysis is restricted to couples in which the man was HIV-positive and the woman was HIV-negative at enrollment and the man was not on antiretroviral treatment. Multivariate Cox models evaluated associations between time-varying contraceptive methods and HIV acquisition among women. Sensitivity analyses explored exposure misclassification and time-varying confounder mediation.

Results—Among 1393 couples, 252 incident infections occurred in women over 2842 couple-years (8.9 infections per 100 couple-years; 95% CI, 7.8–10.0). Multivariate Cox models indicated that neither injectable [adjusted hazard ratio (aHR)=1.2; 95% CI, 0.8–1.7], oral contraceptive pill (OCP, aHR=1.3; 95% CI, 0.9–1.8), or implant (aHR=1.1; 95% CI, 0.5–2.2) use was significantly

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^{*} Corresponding author at: CNR 4011, 1518 Clifton Road NE, Atlanta, GA 30322. Tel.: +1 404 727 9088. kmwall@emory.edu (K.M. Wall)..

associated with HIV acquisition relative to non-hormonal contraception controlling for woman's age, literacy and time-varying measures of genital ulceration/inflammation. This remained true when only looking at the subset of infections acquired from the spouse (82% of infections) and additionally controlling for baseline HIV viral load of the male partner, pregnancy status, and time-varying measures of sperm on a vaginal swab wet prep and self-reported unprotected sex. OCP and injectable users reported more unprotected sex (p<.001), and OCP users were more likely to have sperm on vaginal swab (p=.1) than nonhormonal method users.

Conclusions—We found no association between hormonal contraception and HIV acquisition risk in women. Condom use and reinforced condom counseling should always be recommended for HIV discordant couples. HIV testing of sex partners together is critical to establish HIV risk, ascertain couple fertility intentions and counsel appropriately.

Implications—These findings add to a controversial literature and uniquely address several common design and analytic challenges faced by previous studies. After controlling for confounders, we found no association between hormonal contraception and HIV acquisition risk in women. We support promoting condoms for HIV prevention and increasing the contraceptive method mix to decrease unintended pregnancy.

Keywords

Discordant couples; HIV risk; Hormonal contraception; Longitudinal cohort; Women; Zambia

1. Introduction

Hormonal contraception (HC), including injectable depot medroxyprogesterone acetate (DMPA) and oral contraceptive pills (OCPs), prevents unintended pregnancy [1] and is widely used in high HIV prevalence areas [2]. After reviewing the evidence, a 2014 World Health Organization meeting recommended that no restrictions (Medical Eligibility Criteria Category 1) be placed on HC use by women at risk for HIV and that "women and couples at high risk of HIV acquisition considering progestogen-only injectables should also be informed about and have access to HIV preventive measures, including male and female condoms" [1].

However, extant evidence is conflicting and highly debated [1,3–7]. In a recent systematic review, one out of eight studies deemed "informative but with important limitations" found OCP use significantly increased HIV acquisition risk in women. In that same review, four out of nine studies deemed "informative but with important limitations" found injectable contraception significantly increased HIV acquisition risk in women (notably, one found increased risk in marginal structural models but not Cox models) [8]. In a meta-analysis of observational studies of hormonal contraceptive method use and risk of HIV acquisition among women published in 2015, ten of twelve studies that met inclusion criteria indicated moderate increased risk of HIV acquisition among women using DMPA; none of the 10 studies that met inclusion criteria indicated increased risk among women using OCPs [9]. Finally, in another meta-analysis also published in 2015, DMPA use was associated with HIV acquisition relative to non-hormonal method use after pooling 18 studies, but this association became non-significant when looking at studies deemed to be at lower risk for

methodological bias [10]. Limited data exist evaluating the association between HIV acquisition risk and contraceptive implants [8].

Various analytic design and methodological challenges faced by previous studies have made findings difficult to synthesize. As a result, recommendations were recently developed for more rigorous and consistent analytic methods [11]. The methodological design of the present study allowed for comprehensive consideration of these recommendations. Our objective was to determine the impact of HC, including injectable DMPA, OCPs and implants, on risk of HIV acquisition among women in HIV discordant couples.

2. Methods

2.1. Participants and ethics

From 1994–2012, married or cohabiting HIV sero-discordant (one partner HIV-1 infected and one uninfected) couples living in Lusaka, Zambia were enrolled in a prospective study. Discordant couples were identified from couples' voluntary HIV counseling and testing (CVCT) services offered by the Rwanda Zambia HIV Research Group (RZHRG). RZHRG CVCT promotions, recruitment [12,13], enrollment, retention [14], testing, counseling [14,15] and cohort demographics [16] have been described previously. Briefly, CVCT includes group counseling, rapid HIV testing, and joint post-test couple counseling. This analysis is restricted to couples in which the man was HIV-positive and the woman was HIV-negative (M+F–) at enrollment, the man was not on antiretroviral treatment, and the couple had at least one follow-up visit. This study was approved by the Office for Human Research Protections-registered Institutional Review Boards at Emory University and in Zambia. Written informed consent was obtained from participants.

2.2. Exposure

Contraceptive methods [categorized as no method/condoms alone, combined OCPs (progesterone-only typically prescribed to breastfeeding women until children were 6 months old), DMPA injectables (150 mg IM dosage), copper intrauterine device (IUD), contraceptive implant (Norplant, Jadelle), or permanent methods (hysterectomy/tubal ligation/vasectomy)] were provided at the research site at baseline and at three-monthly follow-up visits. In our primary analysis, contraceptive methods were categorized as implant, injectable, or OCP versus non-HC control (which includes couples using no method/condoms alone, copper IUD, or who had a hysterectomy/tubal ligation/vasectomy).

2.3. Baseline covariates

At enrollment, baseline demographic data were collected including age, years cohabiting, monthly income, and Nyanja literacy (the most commonly used local language in Lusaka). Behavioral risk factors included number of previous pregnancies, current pregnancy, fertility intentions and number of lifetime sexual partners. Clinical characteristics of HIV-positive men partners included viral load (VL) categorized as 100,000 copies/mL, 10,000 to <100,000 copies/mL, or <10,000 copies/mL [17].

2.4. Time-varying covariates

Data collected at scheduled three-monthly follow-up visits included prior three month: incident pregnancy, prevalent pregnancy, self-reported number of protected and unprotected sex acts, any self-reported sex with the study partner with and without a condom and presence of sperm on a vaginal swab wet prep. Composite variables for genital ulceration were created from time-varying measures of chronic/recurrent or acute genital or perianal ulcers (whether diagnosed/treated at the research clinic or reported by the client); ulceration upon physical examination (including erosion or friability of the cervix or vagina in women); or newly positive rapid plasma reagin serology for syphilis [18]. Composite variables for genital inflammation were created from time-varying measures of genital inflammation, genital discharge and trichomoniasis, gonorrhea, chlamydia, candida or bacterial vaginosis [19].

2.5. Outcomes

This analysis considers the association between time-varying HC method use and two outcomes of interest: (1) any incident HIV infection among women partners and (2) incident HIV infection genetically linked to the cohabiting male partner. HIV testing using rapid serologic tests was conducted at three-monthly visits [15]. By comparing conserved PCR-amplified nucleotide sequences from each partner, we determined whether incident infections were genetically linked to the study partner or were unlinked (acquired from outside the study couple) [20]. Eleven couples with unknown linkage were classified as linked [20,21].

2.6. Longitudinal data collection

Data collection varied by type and frequency over 17 years of follow-up. Plasma banking for VL testing was available beginning in 1999. From 1994 to 2002, both partners were seen quarterly, had physical and genital exams, and received laboratory screening for sexually transmitted infections (STIs). Routine p24 antigen screening began in 2003. From 2002 to 2011, fertility goals were recorded. Physical exams and STI laboratory diagnoses were performed at baseline and thereafter given signs and symptoms of STI. In 2007, HIV-negative women were seen at visit months 0, 1, 2 and 3 and completed a sexual exposure risk assessment at quarterly visits. Couples with at least one exposure (unprotected sex, sperm or trichomonas on a wet prep, incident pregnancy or incident STI) received monthly HIV testing until the next quarterly visit, at which time the risk assessment was repeated. From 2008 to 2011, HIV-negative partners were tested monthly.

2.7. Data analysis

Analyses were conducted with SAS v9.3 (Cary, NC, USA). Baseline demographic, behavioral and clinical data are described by HIV transmission status using counts and percentages (for categorical variables) or means and standard deviations (for continuous variables). Index HIV-positive male partner characteristics are only described for genetically linked infections.

Infection rates were calculated as the number of incident infections per couple-year of follow-up, stratified by contraceptive method type. Hormonal method-specific seroincidence

rates were compared to a non-HC reference group using univariate Cox models. Couples were censored if either partner died, the couple separated, the positive partner started antiretroviral therapy or if either partner was lost to follow-up.

Bivariate associations between covariates and outcomes of interest were evaluated via crude hazard ratios (HRs) and 95% confidence intervals (CIs). Multivariate Cox models estimated the effect of time-varying contraceptive method type on incident HIV acquisition. Effect-measure modification was considered for genital ulceration, inflammation, VL of the HIV-positive partner at baseline, fertility intentions, and woman's age. Covariates significantly (p<.05) associated with the exposure and outcome of interest in univariate analyses were considered as potential confounders. Multi-collinearity was assessed; if any two variables were found to be collinear, the variable with the weakest association with the outcome was removed. The proportional hazards assumption was confirmed for time-independent covariates.

2.8. Sensitivity analyses

Sensitivity analyses explored the effects of different contraceptive method exposure categorizations and control groups, misclassification of unprotected sex, controlling for pregnancy, and potential bias due to time-varying confounders simultaneously acting as mediators. To address the first issue, we considered all methods disaggregated versus none/ condom control group; cumulative injectable exposure (calculated as the time-varying cumulative sum of intervals reporting injectable use) versus non-HC control; and cumulative OCP exposure versus non-HC control. To assess misclassification of self-reported unprotected sex, we evaluated the association between this measure and sperm on wet prep in the past three months, incident pregnancy, HIV acquisition, and genital ulceration/ inflammation using Chi-square tests. Models of incident linked seroconversion were run using a composite measure of self-reported unprotected sex (i.e., using an indicator of any self-reported unprotected sex, sperm on vaginal swab wet prep, incident pregnancy, or incident STI). We estimated our results both controlling and not controlling for pregnancy. Marginal structural models estimated through inverse probability weighting were used to adjust for time-varying confounders which may simultaneously act as mediators. Finally, we conducted our analyses among a subset of couples with no indication of condomless sex (using an indicator of any self-reported unprotected sex, sperm on vaginal swab wet prep, incident pregnancy, or incident STI) since the last study interval.

2.9. Unprotected sex, contraceptive method use, and pregnancy

We explored differences in contraceptive method use by measures of unprotected sex using Chi-square tests for categorical variables and t-tests (unequal variance) for continuous variables. We also explored differences in pregnancy status (categorized as pregnant, up to six months post-partum, or not pregnant/post-partum) by measures of protected and unprotected sex using chi-square tests for categorical variables and *t* tests (unequal variance) for continuous variables.

3. Results

3.1. Baseline demographics and bivariate analyses (Tables 1-2)

Eighty-two percent of couples were non-seroconverters (n=1141), 15% acquired genetically linked infections (n=207), and 3% acquired genetically unlinked infections (n=45). Couples were followed for a median of 440 days (IQR=756).

Baseline risk factors significantly associated with incident HIV infection included younger age of the woman, fewer years cohabiting, illiteracy in Nyanja, fewer previous pregnancies, and the woman desiring more children. Additional baseline risk factors significantly associated with incident genetically linked HIV infection included younger age of the man, pregnancy, the man desiring more children, and higher VL of the man (Table 1).

Time-varying covariates significantly associated with incident genetically linked HIV infection included self-reported unprotected sex with the study partner, sperm on a wet prep (Table 2). Time-varying measures of genital inflammation or ulceration in either partner in the past three months were significantly associated with any incident infection and genetically linked infections (data not shown).

3.2. Seroconversion rates by contraceptive method (Table 3)

Of 1393 couples, 252 seroconversions occurred over 2841.9 couple-years (CY) of follow-up. Women using OCPs or injectables since the previous follow-up visit had higher rates of seroconversion relative to women using non-HC methods since the previous follow-up visit; these differences were not statistically significant.

3.3. Multivariate analyses (Table 4)

HC use was not associated in multivariate analyses with any incident HIV infection or the subset of genetically linked infections. No effect-measure modification by genital ulceration, genital inflammation, VL, fertility intentions, or woman's age was observed. Collinear variables included: man and woman age, number of prior pregnancies, and years cohabiting (woman age retained).

Among all infections, use of implant, injectables, or OCPs was not associated with HIV acquisition relative to non-hormonal methods when controlling for woman's age (per year increase), literacy in Nyanja, time-varying measures of genital ulceration and inflammation in the woman partner in the past three months, and time interval since enrollment.

Among linked infections, use of implant, injectables, or OCPs was not associated with HIV acquisition relative to non-hormonal methods controlling for the above factors, baseline pregnancy, sperm present on a vaginal swab wet prep, couples' self-reported unprotected sex in the last three months, time-varying measures of genital ulceration and inflammation of the man in the past three months, and man's baseline log VL.

3.4. Sensitivity analyses

Analyzing different exposure categorizations, controlling for time-varying pregnancy, removing IUD users from the control group, and controlling for fertility intentions did not

yield different conclusions. In almost two-thirds of intervals during which incident HIV was detected, women reported no unprotected sex in the prior three months. Women reported no unprotected sex in almost 40% of intervals during which an incident pregnancy was detected. Using a composite measure to indicate unprotected sex did not yield different results. Marginal structural models did not yield different results (i.e., marginal structural models also did not indicate any association between hormonal contraceptive method use and HIV acquisition risk). Finally, performing these analyses among couples with no indication of condomless sex did not yield different results, and these non-significant findings were of the same magnitude as those for the entire cohort.

3.5. Unprotected sex, contraceptive method use, and pregnancy (Tables 5-6)

OCP users reported a higher number of unprotected sex acts with the study partner in the past three months and had sperm on a wet prep more often than nonhormonal method users. Injectable users reported more unprotected sex than nonhormonal method users. Implant users reported a lower number of unprotected sex acts with the study partner in the past three months, reported less unprotected sex, and had sperm on a wet prep less often than nonhormonal method users (Table 5).

Pregnant women reported a higher average number of protected sex acts relative to post-partum women and a higher average number of unprotected sex acts relative to post-partum or non-pregnant/non-post-partum women. Pregnant women were more likely to report sex without a condom (Table 6).

4. Discussion

Use of oral or injectable HC was not associated with increased risk of HIV acquisition among Zambian women in HIV discordant couples after adjustment for behavioral and biological risk factors. This investigation, both in design and analysis, overcomes several challenges faced by previous studies [11]. We measured various self-reported and biological fixed and time-varying measures of unprotected sex over 17 years of prospective follow-up. We estimated HIV acquisition risk related to contraceptive implants. Contraception was provided at the research site and was measured frequently to capture high rates of stopping and switching; in our cohorts, we have observed that about 25% of women switch methods during follow-up, with most discontinuation/switching observed among injectable (34%) and IUD (33%) users [13,22]. This is one of few studies to differentiate between genetically linked versus unlinked infections, important when modeling index partner covariates. Discordant couples have relatively little within-sample variation in HIV exposure risk. Finally, we corroborated our findings with marginal structural models and rigorous sensitivity analyses.

The relationships between measures of unprotected sex and HC use, pregnancy, and post-partum periods were of particular interest. The reproductive lifetime of most women in Africa cycles through these three stages, each of which involves endogenous and/or exogenous hormonal influences. We found the high rates of biological and self-reported measures of unprotected sex during OCP use, injectable use, and pregnancy surprising. This illustrates the complexity of the relationships between behavioral and biological risk factors

for HIV transmission and indicates that counseling must emphasize maintaining consistent condom use regardless of pregnancy status or other contraceptive method use.

Hormonal implant use was associated with less unprotected sex and fewer pregnancies relative to other methods and had a reduced adjusted hazard ratio for seroconversion. Further research is warranted to assess the role of this effective and cost-effective contraceptive method, along with the copper IUD, in women and couples at risk of HIV. Ongoing long-acting reversible contraception (LARC) promotion and provision for those wishing to delay pregnancy is important given high rates of unprotected sex and unintended pregnancy observed among OCP users – we have previously shown that the rate of unintended pregnancy among OCP users in our discordant couple cohorts (20.7/100 CY) was not significantly lower than women reporting no method/condom use only [23].

Selection bias due to enrollment and loss to follow-up have been thoroughly explored in our cohorts: among M+F-couples, older age and current contraceptive use are predictive of enrollment, while residence far from the clinic, younger age, and women's age at first intercourse being 17 are predictive of attrition [14]. Our findings may therefore be most generalizable to relatively older, contraception experienced couples. Additionally, our study was underpowered to rule out a type II error in conclusions drawn from the univariate associations between seroincidence rates in implant versus non-HC control users.

Based on our findings, we support efforts to increase: 1) the contraceptive method mix to decrease unintended pregnancy, in particular access to LARC methods which are not currently available to many African women, 2) reinforced condom counseling for all persons at risk of HIV, and 3) couple's HIV testing to ascertain the most immediate source of HIV risk of negative adults and support couple-level fertility intentions.

Pragmatically, the latter can be achieved by integrating HIV and family planning services with a focus on couples. Finally, when weighing the current body of published evidence, the effectiveness of HC, especially hormonal LARC methods, to decrease unintended pregnancy, maternal and child mortality, and vertical HIV transmission must be considered when making policy recommendations.

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The corresponding author had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

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References

1. World Health Organization. Hormonal contraceptive methods for women at high risk of HIV and living with HIV: 2014 Guidance statement. Geneva, Switzerland: 2014.

- 2. United Nations Department of Economic and Social Affairs. World Contraceptive Use 2011. 2011
- 3. Westhoff CL, Winikoff B. DMPA and HIV: do we need a trial? Contraception. 2014; 90(4):353. [PubMed: 25183262]
- 4. Jones HE. Time to focus on improving the contraceptive method mix in high HIV prevalence settings and let go of unanswerable questions. Contraception. 2014; 90(4):357–9. [PubMed: 24993486]
- Colvin, CJ.; Harrison, A. Broadening the debate over HIV and hormonal contraception.. Lancet Infect Dis. 2015. http://dx.doi.org/10.1016/S1473-3099(14)71076-X [Epub ahead of print]
- 6. Jain AK. Hormonal contraception and HIV acquisition risk: implications for individual users and public policies. Contraception. 2012; 86(6):645–52. [PubMed: 22541635]
- 7. DMPA and HIV: why we need a trial. Contraception. 2014; 90(4):354-6. [PubMed: 25183263]
- 8. Polis CB, Phillips SJ, Curtis KM, Westreich DJ, Steyn PS, Raymond E, et al. Hormonal contraceptive methods and risk of HIV acquisition in women: a systematic review of epidemiological evidence. Contraception. 2014; 90(4):360–90. [PubMed: 25183264]
- Ralph, LJ.; McCoy, SI.; Shiu, K.; Padian, NS. Lancet. Hormonal contraceptive use and women's risk of HIV acquisition: a meta-analysis of observational studies.. Infect Dis 2015, http://dx.doi.org/ 10.1016/S1473-3099(14)71052-7 [Epub ahead of print]
- Morrison CS, Chen PL, Kwok C, Baeten JM, Brown J, Crook AM, et al. Hormonal contraception and the risk of HIV acquisition: an individual participant data meta-analysis. PLoS Med. 2015; 12(1):e1001778. http://dx.doi.org/10.1371/journal.pmed.1001778. [PubMed: 25612136]
- 11. Polis CB, Westreich D, Balkus JE, Heffron R. Assessing the effect of hormonal contraception on HIV acquisition in observational data: challenges and recommended analytic approaches. AIDS. 2013; 27(Suppl 1):S35–43. [PubMed: 24088682]
- 12. Wall KM, Kilembe W, Nizam A, Vwalika C, Kautzman M, Chomba E, et al. Promotion of couples' voluntary HIV counselling and testing in Lusaka, Zambia by influence network leaders and agents. BMJ Open. 2012; 2(5)
- Mark KE, Meinzen-Derr J, Stephenson R, Haworth A, Ahmed Y, Duncan D, et al. Contraception among HIV concordant and discordant couples in Zambia: a randomized controlled trial. J Womens Health (Larchmt). 2007; 16(8):1200–10. [PubMed: 17937573]
- Kempf MC, Allen S, Zulu I, Kancheya N, Stephenson R, Brill I, et al. Enrollment and retention of HIV discordant couples in Lusaka, Zambia. J Acquir Immune Defic Syndr. 2008; 47(1):116–25.
 [PubMed: 18030162]
- 15. Boeras DI, Luisi N, Karita E, McKinney S, Sharkey T, Keeling M, et al. Indeterminate and discrepant rapid HIV test results in couples' HIV testing and counselling centres in Africa. J Int AIDS Soc. 2011; 14:14–8. http://dx.doi.org/10.1186/1758-2652-14-18. [PubMed: 21439074]
- 16. Stephenson R, Barker J, Cramer R, Hall MA, Karita E, Chomba E, et al. The demographic profile of sero-discordant couples enrolled in clinical research in Rwanda and Zambia. AIDS Care. 2008; 20(3):395–405. [PubMed: 18351489]
- 17. Fideli US, Allen SA, Musonda R, Trask S, Hahn BH, Weiss H, et al. Virologic and immunologic determinants of heterosexual transmission of human immunodeficiency virus type 1 in Africa. AIDS Res Hum Retroviruses. 2001; 17(10):901–10. [PubMed: 11461676]
- 18. Dionne-Odom J, Karita E, Kilembe W, Henderson F, Vwalika B, Bayingana R, et al. Syphilis treatment response among HIV-discordant couples in Zambia and Rwanda. Clin Infect Dis. 2013; 56(12):1829–37. [PubMed: 23487377]
- 19. Song W, He D, Brill I, Malhotra R, Mulenga J, Allen S, et al. Disparate associations of HLA class I markers with HIV-1 acquisition and control of viremia in an African population. PLoS One. 2011; 6(8):e23469. [PubMed: 21858133]
- Trask SA, Derdeyn CA, Fideli U, Chen Y, Meleth S, Kasolo F, et al. Molecular epidemiology of human immunodeficiency virus type 1 transmission in a heterosexual cohort of discordant couples in Zambia. J Virol. 2002; 76(1):397–405. [PubMed: 11739704]

21. Campbell MS, Mullins JI, Hughes JP, Celum C, Wong KG, Raugi DN, et al. Viral linkage in HIV-1 seroconverters and their partners in an HIV-1 prevention clinical trial. PLoS One. 2011; 6(3):e16986. [PubMed: 21399681]

- 22. Haddad L, Wall KM, Vwalika B, Khu NH, Brill I, Kilembe W, et al. Contraceptive discontinuation and switching among couples receiving integrated HIV and family planning services in Lusaka, Zambia. AIDS. 2013; 27(Suppl 1):S93–103. [PubMed: 24088689]
- 23. Wall KM, Haddad L, Vwalika B, Htee Khu N, Brill I, Kilembe W, et al. Unintended pregnancy among HIV positive couples receiving integrated HIV counseling, testing, and family planning services in Zambia. PLoS One. 2013; 8(9):e75353. [PubMed: 24098692]

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Table 1

Descriptive analyses and unadjusted associations between covariates and time to HIV seroconversion (N=1393 M+F- couples)

	Non-sero	Non-seroconverters (N=1141)	Linke	d and u	ınlinked	Linked and unlinked infections (N=252)	ns (N=25	(2)	Linke	d infect	Linked infections (N=207)	=207)		
	Z	%	Z	%	HR	95% CI	1	p (2-tailed)	Z	%	HR	95% CI	I.	p (2-tailed)
Man age (mean, S.D.; HR per year increase)	35.5	7.7							33.5	7.5	0.97	0.95	0.99	<.001
Woman age (mean, S.D.; HR per year increase)	28.8	7.0	26.2	6.3	96.0	0.94	0.98	< .0001	26.3	6.3	96.0	0.94	0.98	<.001
Years cohabiting (mean, S.D.; HR per year increase)	8.3	6.7	6.5	5.6	96.0	0.94	0.98	< .001	6.7	5.7	96.0	0.94	0.99	.002
Monthly family income (mean, S.D.; HR per US dollar increase)	94.0	118.8	62.9	70.4	0.998	0.996	1.000	880.	65.4	74.9	1.00	1.00	1.00	.351
Woman reads Nyanja														
Yes, easily	272	24%	40	16%	ref				35	17%	ref			
With difficulty/not at all	857	76%	203	84%	1.50	1.07	2.11	610.	167	83%	1.45	1.01	2.09	.047
Number of previous pregnancies (mean, S.D.; HR per pregnancy increase)	3.7	2.5	3.1	2.0	0.89	0.84	0.94	<.0001	3.1	2.1	0.90	0.85	96.0	.001
Pregnant at baseline														
Yes	155	14%	47	19%	1.31	96.0	1.80	.094	41	20%	1.43	1.02	2.01	.041
No	986	%98	205	81%	ref				166	%08	ref			
Fertility intentions of man														
Yes, next year	4	12%							18	26%	3.02	1.63	5.61	<.001
Yes, but not next year	107	73%							27	40%	2.04	1.17	3.57	.012
Don't know/No	216	%65							23	34%	ref			
Fertility intentions of woman														
Yes, next year	87	19%	24	28%	2.36	1.41	3.96	.001	20	29%	2.72	1.53	4.86	.001
Yes, but not next year	88	19%	26	30%	1.99	1.21	3.30	.007	21	31%	2.22	1.26	3.94	900.
Don't know/No	278	61%	37	43%	ref				27	40%	ref			
Man lifetime sex partners (mean, S.D.; HR per partner increase)	11.6	16.0							10.4	11.5	1.00	0.98	1.01	.378
Man last year sex partners (mean, S.D.; HR per partner increase)	1.8	1.7							1.8	1.3	0.98	0.91	1.06	.654
Woman lifetime sex partners (per partner increase)	2.8	3.4	2.7	2.6	1.01	0.97	1.05	659.	2.7	2.6	1.00	96.0	1.05	.935
Woman last year sex partners (mean, S.D.; HR per partner increase)	1.0	0.4	1.1	0.5	1.24	0.99	1.55	090.	1.1	0.3	1.07	0.76	1.51	.710
Log viral load of positive partner, log10 copies/mL (mean, S.D.; HR per unit increase)	4.6	1.0							5.1	0.7	1.66	1.36	2.04	<.0001

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Table 2

Unadjusted associations between time-varying covariates and time to HIV seroconversion (N=1393 M+F - couples)

	All couples		Linked and unlinked infections	ınlinked	infectio	sus			Linked infections	tions				
	N intervals	%	N intervals	%	HR	95% CI		p (2-tailed)	N intervals	%	HR	95% CI	CI	p (2-tailed)
Contraceptive method used at visit														
* Non-hormonal	0906	%59	153	61%	ref				133	64%	ref			
Implant	1000	%/	6	4%	1.16	0.57	2.35	.678	9	3%	1.01	0.43	2.36	.991
Injectables	1899	14%	41	16%	1.21	0.85	1.73	.296	33	16%	1.15	0.77	1.70	.505
OCPs	1921	14%	49	19%	1.31	0.94	1.82	.114	35	17%	1.10	0.75	1.62	.612
Pregnant during interval														
Yes	1130	%6	29	12%	1.27	98.0	1.88	.225	27	14%	1.41	0.94	2.12	760.
No	11168	91%	213	%88	ref				171	%98	ref			
No. times sex with study partner with a condom in the last 3 months (mean, S.D.)	18.9	19.4	19.4	25.2	1.00	0.99	1.01	.921	19.9	26.5	1.001	1.00	1.01	.789
No. times sex with study partner without a condom in the last 3 months (mean, S.D.)	2.4	8.5	3.6	10.7	1.00	0.99	1.01	.987	3.7	10.3	1.00	0.99	1.01	.890
Sex with study partner with a condom in past 3 months														
Yes	11606	%98							176	85%	1.20	0.81	1.77	0.360
No	1913	14%							30	15%	ref			
Sex with study partner without a condom in past 3 months														
Yes	4087	30%							87	42%	1.39	1.04	1.84	0.024
No	9433	%0/							119	28%	ref			
Sperm present on wet prep														
Yes	831	%9	26	11%	1.50	0.97	2.32	990.	25	13%	1.74	1.11	2.73	.016
No	12209	94%	208	%68	ref				165	81%	ref			

 $^{^{\}ast}$ IUD, none/condoms alone, permanent method.

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 $\label{eq:Table 3} \textbf{Seroconversion rates among HIV-negative women in discordant relationships by method of contraception $(N=1393 M+F-couples)$$

	Number of seroconversions	Couple-years of follow-up time	Seroincidence per 100 couple-years	95%	CI	p (2-tailed)*
Current contraceptive method used at follow-up visit						
Non-hormonal ^	153	1902.3	8.0	6.8	9.4	ref
OCPs	49	424.5	11.5	8.5	15.3	.114
Injectables	41	392.3	10.5	7.5	14.2	.296
Implant	9	122.9	7.3	3.3	13.9	.678
Total	252	2841.9	8.9	7.8	10.0	

 $^{67\} study$ intervals (accounting for $5.7\ couple$ years) were missing contraceptive information.

[^]Copper intrauterine device, none/condoms alone, permanent method.

^{*}From univariate Cox proportional hazards models.

Table 4

Multivariate models of hormonal contraception use and time to HIV seroconversion (N=1393 M+F – couples)

	Linked	and un	linked	infections	Linked i	nfection	ns	
	aHR*	95%	CI	p value (2-tailed)	aHR**	95%	CI	p value (2-tailed)
Current contraceptive method at follow-up visit								
Non-hormonal ^	ref				ref			
Implant	1.08	0.53	2.20	.83	0.96	0.29	3.14	.947
Injectables	1.19	0.81	1.73	.37	1.34	0.85	2.12	.204
OCPs	1.29	0.92	1.80	.15	1.39	0.90	2.15	.140

aHR: adjusted hazard ratio.

^{*} Controlling for woman's age (per year increase), woman's literacy in Nyanja, genital ulceration of woman in past 3 months, genital inflammation of woman in the past 3 months, and time interval since enrollment (0-3 months versus > 3 months).

^{**} Controlling for * and baseline pregnancy, sperm present on a wet prep, couples' self-reported unprotected sex in the last 3 months, genital ulceration of man in past 3 months, genital inflammation of man in the past 3 months, and man's baseline log viral load (per log10 copies/mL increase)

[^]Copper intrauterine device, none/condoms alone, permanent method.

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 $\label{eq:Table 5} \textbf{Measures of unprotected sex by method of contraception (N=1393 M+F-couples)}$

	Non-hormonal (condoms alone, method)		OCPs		Injectable		Implant		p (2-tailed)*
	N intervals	%	N intervals	%	N intervals	%	N intervals	%	
Number of times sex with partner in project without a condom in the last 3 months reported by woman (mean, S.D.)	2.47	8.78	2.87	9.10	2.42	7.96	1.25	4.88	^&
Sex with study partner without a condom in past 3 months reported by woman									^#&
Yes	2644	29%	717	37%	640	34%	180	18%	
No	6321	71%	1198	63%	1221	66%	810	82%	
Sperm present on wet prep in last 3 months									
Yes	570	7%	151	8%	113	6%	23	2%	
No	7836	93%	1774	92%	1805	94%	986	98%	

^{*} chi-Square (or Fisher's exact) test for categorical variables; t-tests (unequal variance) for continuous variables.

p<.1 for tests of differences between OCP versus non-hormonal contraception distributions.

 $^{^{\#}}$ p<.001 for tests of differences between injectables versus non-hormonal contraception distributions.

 $^{^{\&}amp;}$ p<.001 for tests of differences between implant versus non-hormonal contraception distributions.

 $\label{eq:Table 6} \textbf{Measures of protected and unprotected sex by pregnancy status (N=1393 M+F-couples)}$

	Pregnant		Post-partum (u past delivery)	p to 6 months	Not pregnant o	or post-	p (2-tailed)*
	N intervals	%	N intervals	%	N intervals	%	
Number of times sex with partner in project with a condom in the last 3 months reported by woman (mean, S.D.)	16.67	17.09	8.77	12.05	19.59	20.09	^#&
Number of times sex with partner in project without a condom in the last 3 months reported by woman (mean, S.D.)	6.10	14.42	1.53	6.59	2.25	8.05	^#&
Sex with study partner with a condom in past 3 months reported by woman							^#&
Yes	959	84%	322	63%	9257	87%	
No	180	16%	187	37%	1377	13%	
Sex with study partner without a condom in past 3 months reported by woman							
Yes	593	52%	116	23%	3145	30%	^#&
No	546	48%	393	77%	7490	70%	
Sperm present on wet prep in last 3 months							#
Yes	95	9%	32	6%	690	7%	
No	966	91%	478	94%	9476	93%	

^{*} chi-Square (or Fisher's exact) test for categorical variables; t tests (unequal variance) for continuous variables.

 $^{\ ^{\}wedge}$ p<.01 for tests of differences between pregnant versus post-partum women.

[#]p<.01 for tests of differences between pregnant versus not pregnant or post-partum women.

 $[\]ensuremath{\&}_{p<.01}$ for tests of differences between post-partum versus not pregnant or post-partum women.